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- (54) Title: WATER-SOFTENING AND DETERGENT COMPOSITIONS
- (57) Abstract

Tablets containing a water-softening agent, which may be tablets of detergent composition for fabric washing, contain a crystalline acetate or citrate to promote disintegration at the time of use. To inhibit caking and facilitate handling during manufacture, smaller particles of another substance are provided at the surface of the crystals of the acetate or citrate.

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# WATER-SOFTENING AND DETERGENT COMPOSITIONS

This invention relates to compositions in the form of tablets, containing a water-softening agent. These tablets may be embodied as detergent compositions for use in fabric washing, or as water-softening tablets, which could be used in fabric washing jointly with a composition containing detergent active, or could possibly be used in other applications, e.g. in machine dishwashing as an anti-limescale product.

Detergent compositions in tablet form are described, for example, in GB 911204 (Unilever), US 3953350 (Kao), JP 60-15 015500A (Lion), JP 60-135497A (Lion) and JP 60-135498A (Lion); and are sold commercially. Tablets have several advantages over powdered products: they do not require measuring and are thus easier to handle and dispense into the washload, and they are more compact, hence facilitating more economical storage.

Detergent tablets are generally made by compressing or compacting a detergent powder, which includes detergent active and detergency builder. EP-A-522766 explains that difficulty has been found in providing tablets which have adequate strength when dry, yet disperse and dissolve quickly when added to wash water. The problem has proved especially difficult with compositions containing

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insoluble aluminosilicate as detergency builder but the problem also arises with tablets which contain sodium tripolyphosphate as the detergency builder.

This prior document teaches that at least some particles of the composition should be coated with a binder which helps to hold the tablet together and allows a tablet to be made using a lower compaction pressure. The binder can also function as a disintegrant.

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EP-A-482627 teaches that a detergent composition for compaction into tablets with improved solubility should include potassium carbonate together with nonionic surfactant.

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EP-A-711827 teaches that speed of disintegration of tablets can be improved by including a highly water-soluble citrate. Tablet compositions exemplified in that document include sodium citrate dihydrate and also polyethylene glycol as an organic polymeric binder. This document also mentions that sodium acetate can be included in a composition as a lubricant to aid tabletting. The trihydrate of sodium acetate is not named. The amount of lubricant is not stated, but it would be appropriate to include only a small amount.

WO 90/02165 mentions a range of materials including sodium acetate trihydrate as tableting aids, preferably used as a

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small percentage of the composition and preferably of fine particle size. A range of possible functions is attributed to these tableting aids.

Our European patent application 97308427.0 now published on 29 April 1998 as EP-A-838519 discloses that a tablet of a compacted particulate composition which contains a water-softening agent can be made to disintegrate more rapidly at the time of use by incorporating into the tablet, mixed with the water-softening agent, a substance selected from sodium acetate trihydrate, potassium acetate and mixtures thereof.

We have now found, however, that when these disintegrant

salts are handled on a commercial scale, they have a

tendency to cake into inconvenient lumps even though they

are simple crystalline solids. We have found that this

problem, which we believe has not previously been

recognised, can be reduced by applying finely divided

particulate material to the exterior of the crystals.

Moreover, the benefit of improved speed of disintegration
is substantially retained.

Accordingly, in a first aspect, this invention provides a

25 process for the production of a tablet of a compacted

particulate composition by mixing

- (i) a water-softening agent, and
- (ii) a water-soluble crystalline salt selected from sodium

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citrate dihydrate, sodium acetate trihydrate,
potassium acetate and mixtures thereof
and compacting the resulting mixed composition into
tablets or regions of tablets, characterised by the
presence of particles of another substance at the surface
of the crystals of the said crystalline salt (ii) before
it is mixed with the water softening agent (i).

In a second aspect this invention provides a tablet of

compacted particulate composition containing a watersoftening agent mixed with a crystalline salt selected
from sodium citrate dihydrate, sodium acetate trihydrate,
potassium acetate and mixtures thereof characterised by
particles of another material at the surface of the

crystals of the said crystalline salt.

The process may include a step of application of particles of material to the surface of crystals of the crystalline salt. However, this may be carried out by the manufacturer of that salt, at the place and time of its production, prior to transport to the place where the tablets are made by mixing and compaction.

In the tablets of the present invention and likewise in the particulate compositions made by mixing the ingredient materials (i), (ii) and optionally other materials (iii) preparatory for compaction into tablets, the amount of water-softening agent will generally be at least 15% by

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weight of the composition. Depending on the function for which the tablets are intended, the amount may range up to 90 or 93% by weight. In significant forms of this invention there is at least 15%, by weight of the composition, of a water-insoluble water softening agent.

The amount of the ingredient (ii) ie sodium citrate dihydrate, sodium acetate trihydrate, potassium acetate or mixture of them, may be at least 7% by weight of the composition, often at least 10% or 13% by weight. It will generally not exceed 35% by weight of the composition and frequently will not exceed 25% or 30% by weight of the composition.

In another aspect, this invention provides the use of crystals of sodium acetate trihydrate, potassium acetate or mixture of them, bearing particles of another substance at the surface of the crystals of the said salt, in a tablet of compacted particulate composition or a region thereof, to enhance the disintegration of the tablet in water.

Sodium acetate trihydrate and potassium acetate are preferred over sodium citrate dihydrate. Although potassium acetate is very effective, it is hygroscopic. We have found it easier to use sodium acetate trihydrate which is therefore the material of preference. If a mixture of these materials is used, it is preferred that

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sodium acetate trihydrate provides at least 5% by weight of the composition which is compacted into a tablet or region of a tablet.

- It is strongly preferred that the crystals of sodium citrate dihydrate, sodium acetate trihydrate and/or potassium acetate have a mean particle size of above 250μm, preferably above 300μm (0.3mm), better above 500μm (0.5mm) to facilitate handling prior to and during compaction. The crystal size will probably have a mean value less than 2mm, preferably less than 1mm. The amount of such particles is preferably at least 7%, better at least 10% or 13% of the weight of the composition.
- Suitably, the material at the surface of the crystals has a smaller particle size than the crystals. The mean particle size of this material may be no more than  $180\mu m$  or  $100\mu m$ . With some materials the mean particle size may be no more than  $20\mu m$  and it may be no more than  $10\mu m$  or  $5\mu m$ , especially if it is water-insoluble. Thus the material on the surface of the crystals may have a mean particle size which is not more than one tenth or one thirtieth the mean size of the crystals.
- A number of substances have been found suitable for application to the surface of particles of the crystalline salt. Materials which have found to be suitable include alkali metal carbonate and bicarbonates, sodium

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aluminosilicates and particles of polyethylene glycol.

Particles of sodium aluminosilicate are particularly preferred because they function as a water-softening agent when the composition is used.

#### Water-softening agent

- It is particularly envisaged that this invention will be applied to tablets containing water-insoluble water softening agent, notably alkali-metal aluminosilicate.

  However, it could be applied in tablets containing a soluble water-softening agent such as a condensed phosphate. It could be applied in tablets containing both soluble and insoluble water softening agents as might be used in countries where a restricted quantity of phosphate detergency builder is permitted.
- It is very well known that water-insoluble alkali metal aluminosilicates can function to soften water, removing calcium ions and to a lesser extent magnesium ions by ion exchange. Aluminosilicates have become strongly favoured as environmentally acceptable detergency builders.

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Alkali metal (preferably sodium) aluminosilicates used in tablets of the present invention may be either crystalline, amorphous or a mixture of the two. Such

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aluminosilicates generally have a calcium ion exchange capacity of at least 50 mg CaO per gram of aluminosilicate, comply with a general formula:

0.8-1.5  $Na_2O$  .  $Al_2O_3$  . 0.8-6  $SiO_2$ 

and incorporate some water. Preferred sodium
aluminosilicates within the above formula contain 1.5-3.5
SiO<sub>2</sub> units. Both amorphous and crystalline
aluminosilicates can be prepared by reaction between
sodium silicate and sodium aluminate, as amply described
in the literature.

Suitable crystalline sodium aluminosilicate ion-exchange detergency builders are described, for example, in GB 1429143 (Procter & Gamble). The preferred sodium aluminosilicates of this type are the well known commercially available zeolites A and X, and mixtures thereof. Also of interest is the novel zeolite P described and claimed in EP 384070 (Unilever).

Another category of water-insoluble material which can function as a water-softening agent and detergency builder is the layered sodium silicate builders disclosed in US-A-4464839 and US-A-4820439 and also referred to in EP-A-551375.

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These materials are defined in US-A-4820439 as being crystalline layered sodium silicate of the general formula  ${\rm NaMSi}_{\rm x}{\rm 0}_{2{\rm x}+1}~.~{\rm YH}_2{\rm O}$ 

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where M denotes sodium or hydrogen,
x is from 1.9 to 4 and y is from 0 to 20.

Quoted literature references describing the preparation of such materials include Glastechn. Ber. 37, 194-200 (1964), Zeitschrift für Kristallogr. 129, 396-404 (1969), Bull. Soc. Franc. Min. Crist., 95, 371-382 (1972) and Amer. Mineral, 62, 763-771 (1977). These materials also function to remove calcium and magnesium ions from water.

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It is customary to use a water-soluble builder (water-softening agent) jointly with aluminosilicate, to enhance water-softening efficacy. Such water-soluble co-builders are generally used in an amount which is not greater than the amount of aluminosilicate, often less than half the amount of aluminosilicate. Water-soluble builders may be organic or inorganic. Inorganic builders that may be present include alkali metal (generally sodium) carbonate; while organic builders include polycarboxylate polymers, such as polyacrylates, acrylic/maleic copolymers, and acrylic phosphonates, monomeric polycarboxylates such as citrates, gluconates, oxydisuccinates, glycerol mono- diand trisuccinates, carboxymethyloxysuccinates, carboxymethyloxymalonates and hydroxyethyliminodiacetates.

Especially preferred supplementary builders are polycarboxylate polymers, more especially polyacrylates

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and acrylic/maleic copolymers, and monomeric polycarboxylates, more especially citric acid and its salts.

If a tablet contains only soluble water-softening agent, this may well be sodium tripolyphosphate, which is widely used as a detergency builder in some countries.

When using aluminosilicate or other insoluble detergency builder/water-softening agent it is often a commercial or legislative requirement to avoid phosphates. Some tablet compositions of the invention do not contain more than 5 wt% of inorganic phosphate builders, and are desirably substantially free of phosphate builders. However,

tableted compositions containing some phosphate builder are also within the broad scope of the invention. In particular, a tablet or region thereof may contain at least 15 wt% insoluble water softening agent, with phosphate or other water-soluble builder in addition.

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As mentioned above, compositions of this invention may be embodied as detergent compositions for use in fabric washing, in which case the composition will generally contain from 15 to 60% by weight of detergency builder, notably water-insoluble aluminosilicate, together with 5 to 50% by weight of one or more detergent-active compounds. Such a composition may well contain from 0.5 to 15% by weight of a supplementary builder, notably

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polycarboxylate, and also other detergency ingredients.

Another possibility is that the invention may be embodied in tablets whose principal or sole function is that of removing water hardness. In such tablets the watersoftening agents, especially water-insoluble aluminosilicate, may provide from 50 to 98% of the tablet composition. A water-soluble supplementary builder may well be included, for instance in an amount from 2% to 30wt% of the composition.

Water-softening tablets embodying this invention may include some detergent active. Notably, water-softening tablets may include nonionic surfactant which can act as a lubricant during tablet manufacture and as a low foaming detergent during use. The amount may be small, e.g. from 0.2 or 0.5% by weight of the composition up to 3% or 5% by weight.

#### 20 <u>Detergent Tablets</u>

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Tablets for use in fabric washing will generally contain from 5% to 50% by weight of detergent active, preferably from 5% or 9wt% up to 40% or 50wt%. Detergent-active material present may be anionic (soap or non-soap), cationic, zwitterionic, amphoteric, nonionic or any combination of these.

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Anionic detergent-active compounds may be present in an amount of from 0.5 to 40 wt%, preferably from 2% or 4% to 30% or 40wt%.

5 Synthetic (i.e. non-soap) anionic surfactants are well known to those skilled in the art. Examples include alkylbenzene sulphonates, particularly sodium linear alkylbenzene sulphonates having an alkyl chain length of  $C_8$ - $C_{15}$ ; olefin sulphonates; alkane sulphonates; dialkyl sulphosuccinates; and fatty acid ester sulphonates.

Primary alkyl sulphate having the formula  $ROSO_3^- M^+$ 

in which R is an alkyl or alkenyl chain of 8 to 18 carbon atoms especially 10 to 14 carbon atoms and M<sup>+</sup> is a solubilising cation, is commercially significant as an anionic detergent active. It is frequently the desired anionic detergent and may provide 75 to 100% of any anionic non-soap detergent in the composition.

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In some forms of this invention the amount of non-soap anionic detergent lies in a range from 0.5 to 15 wt% of the tablet composition.

It may also be desirable to include one or more soaps of fatty acids. These are preferably sodium soaps derived from naturally occurring fatty acids, for example, the fatty acids from coconut oil, beef tallow, sunflower or

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hardened rapeseed oil.

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Suitable nonionic detergent compounds which may be used include in particular the reaction products of compounds having a hydrophobic group and a reactive hydrogen atom, for example, aliphatic alcohols, acids, amides or alkyl phenols with alkylene oxides, especially ethylene oxide either alone or with propylene oxide.

- 10 Specific nonionic detergent compounds are alkyl  $(C_{8-22})$  phenol-ethylene oxide condensates, the condensation products of linear or branched aliphatic  $C_{8-20}$  primary or secondary alcohols with ethylene oxide, and products made by condensation of ethylene oxide with the reaction
- products of propylene oxide and ethylene-diamine. Other nonionic detergent compounds include alkylpolyglycosides, long-chain amine oxides, tertiary phosphine oxides, and dialkyl sulphoxides.
- 20 Especially preferred are the primary and secondary alcohol ethoxylates, especially the  $C_{9-11}$  and  $C_{12-15}$  primary and secondary alcohols ethoxylated with an average of from 5 to 20 moles of ethylene oxide per mole of alcohol.
- In certain forms of this invention the amount of nonionic detergent lies in a range from 4 to 40%, better 4 or 5 to 30% by weight of the composition.

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Many nonionic detergent-active compounds are liquids.

These may be absorbed on a porous carrier. Preferred carriers include zeolite; zeolite granuled with other materials, for example Wessalith CS (Trade Mark),

Wessalith CD (Trade Mark) or Vegabond GB (Trade Mark); sodium perborate monohydrate; Burkeite (spray-dried sodium carbonate and sodium sulphate as disclosed in EP-A-221776 of Unilever); and layered sodium silicate as described in US-A-4664839.

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#### Bleach System

Tableted detergent compositions according to the invention may contain a bleach system. This preferably comprises

15 one or more peroxy bleach compounds, for example, inorganic persalts or organic peroxyacids, which may be employed in conjunction with activators to improve bleaching action at low wash temperatures. If any peroxygen compound is present, the amount is likely to lie in a range from 10 to 25% by weight of the composition.

Preferred inorganic persalts are sodium perborate monohydrate and tetrahydrate, and sodium percarbonate, advantageously employed together with an activator. Bleach activators, also referred to as bleach precursors, have been widely disclosed in the art. Preferred examples include peracetic acid precursors, for example, tetraacetylethylene diamine (TAED), now in widespread

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commercial use in conjunction with sodium perborate; and perbenzoic acid precursors. The quaternary ammonium and phosphonium bleach activators disclosed in US 4751015 and US 4818426 (Lever Brothers Company) are also of interest.

Another type of bleach activator which may be used, but which is not a bleach precursor, is a transition metal catalyst as disclosed in EP-A-458397, EP-A-458398 and EP-A-549272. A bleach system may also include a bleach stabiliser (heavy metal sequestrant) such as ethylenediamine tetramethylene phosphonate and

diethylenetriamine pentamethylene phosphonate.

As indicated above, if a bleach is present and is a water-soluble inorganic peroxygen bleach, the amount may well be from 10% to 25% by weight of the composition.

### Other Ingredients

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Detergent tablets of the invention may also contain one of
the detergency enzymes well known in the art for their
ability to degrade and aid in the removal of various soils
and stains. Suitable enzymes include the various
proteases, cellulases, lipases, amylases, and mixtures
thereof, which are designed to remove a variety of soils
and stains from fabrics. Examples of suitable proteases
are Maxatase (Trade Mark), as supplied by Gist-Brocades
N.V., Delft, Holland, and Alcalase (Trade Mark), and
Savinase (Trade Mark), as supplied by Novo Industri A/S,

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Copenhagen, Denmark. Detergency enzymes are commonly employed in the form of granules or marumes, optionally with a protective coating, in amount of from about 0.1% to about 3.0% by weight of the composition; and these granules or marumes present no problems with respect to compaction to form a tablet.

The detergent tablets of the invention may also contain a fluorescer (optical brightener), for example, Tinopal (Trade Mark) DMS or Tinopal CBS available from Ciba-Geigy AG, Basel, Switzerland. Tinopal DMS is disodium 4,4'bis-(2-morpholino-4-anilino-s-triazin-6-ylamino) stilbene disulphonate; and Tinopal CBS is disodium 2,2'-bis-(phenyl-styryl) disulphonate.

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An antifoam material is advantageously included, especially if the detergent tablet is primarily intended for use in front-loading drum-type automatic washing machines. Suitable antifoam materials are usually in granular form, such as those described in EP 266863A (Unilever). Such antifoam granules typically comprise a mixture of silicone oil, petroleum jelly, hydrophobic silica and alkyl phosphate as antifoam active material, sorbed onto a porous absorbed water-soluble carbonate-based inorganic carrier material. Antifoam granules may be present in an amount up to 5% by weight of the composition.

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It may also be desirable that a detergent tablet of the invention includes an amount of an alkali metal silicate, particularly sodium ortho-, meta- or preferably alkali metal silicates at levels, for example, of 0.1 to 10 wt%, may be advantageous in providing protection against the corrosion of metal parts in washing machines, besides providing some measure of building and giving processing benefits.

- Further ingredients which can optionally be employed in the detergent tablet of the invention include antiredeposition agents such as sodium carboxymethylcellulose, straight-chain polyvinyl pyrrolidone and the cellulose ethers such as methyl cellulose and ethyl hydroxyethyl cellulose, fabric-softening agents; heavy metal sequestrants such as EDTA; perfumes; colourants or coloured speckles, and tabletting aids such as binders and lubricants.
- The particulate mixed composition which is compacted into tablets may in principle have any bulk density. However, the present invention is especially relevant to tablets made by compacting powders of relatively high bulk density, because of their greater tendency to exhibit disintegration and dispersion problems. Such tablets have the advantage that, as compared with a tablet derived from a low bulk density powder, a given dose of composition can be presented as a smaller tablet.

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Thus the starting particulate composition may suitably have a bulk density of at least 400 g/litre, preferably at least 500 g/litre, and advantageously at least 700 g/litre.

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A tablet of the invention may be either homogeneous or heterogeneous. In the present specification, the term "homogeneous" is used to mean a tablet produced by compaction of a single particulate composition, but does not imply that all the particles of that composition will necessarily be of identical composition. Indeed it is likely that the composition will contain the sodium acetate trihydrate or potassium acetate as separate particles.

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The term "heterogeneous" is used to mean a tablet consisting of a plurality of discrete regions, for example layers, inserts or coatings, each derived by compaction from a particulate composition and large enough to constitute from 10 to 90% of the weight of the whole tablet.

It is possible that the potassium acetate, sodium acetate trihydrate or sodium citrate dihydrate will be contained within one or more but not all such discrete regions of a heterogeneous tablet, such as a layer or an insert. The presence of such a layer or insert could assist break up of the entire tablet when placed in water.

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Preferably, the composition of the tablet or a tablet region contains particles in which detergent active is mixed with other materials, and separate particles of the crystalline salt, especially sodium acetate trihydrate, desirably having a mean particle size over 0.3mm. Thus the water-softening agent which is ingredient (i) may be granulated with detergent and then mixed with the crystalline salt (ii) and optional other ingredients (iii) to provide the mixture which is compacted into tablets or regions of tablets. Particles of the crystalline salt, separate from the detergent active, may be at least 7%, better at least 10% or 13% by weight of the composition and the tablet or tablet region compacted therefrom.

#### 15 <u>Tableting</u>

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Tableting entails compaction of a particulate composition.

A variety of tableting machinery is known, and can be used. Generally it will function by stamping a quantity of the particulate composition which is confined in a die.

Tableting may be carried out at ambient temperature or at a temperature above ambient which may allow adequate strength to be achieved with less applied pressure during compaction. In order to carry out the tableting at a temperature which is above ambient, the particulate composition is preferably supplied to the tableting machinery at an elevated temperature. This will of course

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supply heat to the tableting machinery, but the machinery may be heated in some other way also.

If any heat is supplied, it is envisaged that this will be supplied conventionally, such as by passing the particulate composition through an oven, rather than by any application of microwave energy. However, this invention could be utilised in a process in which the tableting step includes application of microwave energy to the composition.

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### Example 1

Sodium carbonate and bicarbonate were demonstrated to reduce caking of sodium acetate trihydrate, using the following test procedure:

Crystalline sodium acetate trihydrate (supplied by Verdugt) with average particle size  $770\mu m$  was mixed with sodium carbonate or sodium bicarbonate in varying amounts up to 5% by weight.

The sodium carbonate was light soda ash (supplied by Akzo). It was anhydrous and had an average particle size below 200  $\mu\text{m}$ , estimated as 140  $\mu\text{m}$ .

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The sodium bicarbonate (supplied by Solvay) was likewise anhydrous and was passed through a  $180\,\mu\text{m}$  sieve before use. The average particle size of the sieved material was estimated to be about  $90\,\mu\text{m}$ .

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3.5kg quantities of sodium acetate trihydrate were mixed by hand with the sodium carbonate or bicarbonate. Any lumps present in the sodium acetate trihydrate were removed and broken up or discarded prior to weighing out the 3.5kg quantity.

After mixing, the mixture was stored in a closed bucket for various periods at 20°C or 37°C. Before and after

storage a portion of the sodium acetate was poured through a sieve with 3.35mm apertures. Material retained on the sieve was considered caked. It was weighed and expressed as a percentage of the whole quantity. The following results were obtained:

% caked after storage period Additive Temp before 1 33 (°C) day storage days days days days none 20 15.4 22.0 30.4 51.0 37 0 27.9 24.5 47.1 54 73.3 2.5% 20 0 11.1 25.5 23.8 24.5 34.8 carbonate 37 57.3 0 22.9 36.5 44.1 5% 0 6.4 15.7 9.5 14.6 29.9 20 carbonate 0 22.3 27.0 36.4 40.5 37

% caked after storage period 10 days 43 days Additive Temp before 41 days (°C) storage 0.75 2% 20 0 1.4 bicarbonate 48.1 37 0

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# Example 2

The previous example was repeated with further materials all of which were inorganic, as follows:

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Alusil N, a commercial aluminosilicate flow aid available from Crosfields, mean particle size  $6\mu m$ .

Zeolite 4A, mean particle size in a range from 2 to  $5\mu m$ .

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Zeolite A24, a maximum aluminium zeolite P available from Crosfields, mean particle size in a range from 0.7 to  $1.5\mu\text{m}$ 

15 Storage was for seven days in every case.

Material passing through the 3.35mm sieve was tested for its stickiness by the following procedure referred to as "compression test". A cylindrical mould made in two

20 halves is placed on a flat surface with its axis vertical. It then defines a cylindrical chamber 9cm in diameter and 11cm high. This is filled with the material to test. The material is next compressed within the mould by means of a 10kg weight for two minutes. The weight and the mould are then removed to leave a free-standing cylindrical compact of the test material. Weight is progressively applied to the top of this compact until collapse. The result is expressed as the applied weight in grams.

The following results were obtained:

	બ	caked		compress	sion tes	t (gm)
Additive	before storage	7 days at 20°C	7 days at 37°C	before storage	7 days at 20°C	7 days at 37°C
none	0%	40.9%	64.6%	952	707	1206
0.2% A24	0%	0.6%	7.9%	2457	1959	2458
0.4% A24	0%	1.0%	7.7%	2457	2457	2458
0.6% A24	0%	0.7%	3.7%	2457	2457	2959
1% A24	0 %	0.1%	0.35%	2457	2208	2208
2% A24	0%	0.3%	0.4%	2958	1708	2208
0.6% 4A	0%	11.6%	18.5%	2209	1959	1959
0.6% Alusil-N	0%	0.9%	1.3%	1708	1457	1707

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In can be seen from the results in this table that the application of these materials increases the stickiness of the material compared to sodium acetate trihydrate alone.

In spite of this however, the caking into lumps is dramatically reduced.

## Example 3

Example 2 was repeated, using as additive polyethylene glycol of molecular weight 1500. This was in the form of fine powder which was passed through a 180 µm sieve before use. Its mean particle size was estimated as about  $90\mu m$ .

The following results were obtained:

10		% caked af	ter	compression	n test
	% PEG	8 days at 20°C	8 days at 37°C	8 days at 20°C	8 days at 37°C
	none	40.8%	60.9%	1099	1200
	0.5%	31.3%	48.6%	702	350
	1%	26.4%	44.8%	601	350
15	2%	33.5%	37.8%	601	450
	4%	22.4%	41.2%	low	700

It can be seen from the above table that the PEG 1500 was effective to reduce caking. Moreover, it was observed 20 that the lumps which were formed were relatively soft and easily broken whereas lumps formed when the sodium acetate trihydrate was not treated with polyethylene glycol were harder lumps. This difference is consistent with the compression test results where it can be seen that the application of polyethylene glycol reduced the stickiness 25 of sodium acetate trihydrate.

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#### Example 4

Tablets suitable for use in water-softening were made from mixtures of zeolite granules and sodium acetate trihydrate with zeolite particles on the surface of the sodium acetate trihydrate crystals.

The zeolite granules were a commercial product available from Norsohaas under designation WLZ-10. Their composition was:

Polycarboxylate 9-11%

Zeolite A 69.5-73.5%

Water 17-20%

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The polycarboxylate was a copolymer of acrylate and maleate. Such polymers are known as water-soluble builders which enhance the water-softening efficacy of zeolite and also inhibit redeposition of soil from a wash liquor. In these granules the polycarboxylate serves as a binder for the zeolite powder.

The granulometry of WLZ-10 was determined as:

25 Rosin Rammler average particle size 625 microns

Rosin Rammler N value 1.88

Bulk density 777 kg/m<sup>3</sup>

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The sodium acetate trihydrate was a technical grade from Verdugt having average particle size  $770\mu m$  and containing 5% of fines, smaller than  $180\mu$ . The sodium acetate trihydrate was mixed with zeolite A24 as used in Example 2 in a quantity of 0.6% based on the weight of sodium acetate trihydrate.

The WLZ-10 zeolite granules and the sodium acetate trihydrate, with zeolite on its surface, were dry mixed in 3:1 weight ratio and then portions of each mixture were stamped into tablets.

## Example 5

Sodium acetate trihydrate (from Verdugt, mean particle size 770μm) was mixed with 2% of its own weight of polyethylene glycol of mean molecular weight 1500 (PEG 1500) in the form of fine powder. This sodium acetate trihydrate plus PEG 1500 mixture was subsequently mixed with a granulated base powder and other ingredients as set out in the following tables. As a comparison sodium acetate trihydrate was used without admixed PEG 1500. This comparative formulation is also shown in the following tables.

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Granulated Base Powder	Parts by weight
Linear alkylbenzene sulphonate	9.4
Nonionic detergent	4.1
Sodium carbonate	3.1
Soap	0.7
Sodium carboxymethyl cellulose	0.4
Zeolite A24 (anhydrous)	20.9
Sodium acetate trihydrate	2.7
Moisture and non-detergent organic material	3.7
TOTAL	45

		% k	y weight
		with PEG	comparative
15	Base powder	45	45
	Sodium percarbonate	15.3	15.3
	TAED (83% active) granules	5.2	5.2
	Na-disilicate (80% silicate)	3.6	3.6
	Phosphonate sequestrant	0.7	0.7
20	Soil release polymer	1.1	1.1
	Antifoam granules (18% active)	1.8	1.8
	Fluorescer granules (15% active)	1.0	1.0
	Acrylate maleate copolymer	1.3	1.3
	Sodium carbonate	2.0	2.0
25	Sodium acetate trihydrate + 2% PEG 1500	23.0	-
	Sodium acetate trihydrate	_	23.0
	TOTAL	100	100

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Tablets were made from these two formulations, using A Carver laboratory press to make cylindrical tablets with a

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weight of 35 gm. Various amounts of force were used to stamp the tablets.

The resulting tablets were tested by the following test for tablet strength. A tablet is placed between the platens of a materials testing machine so that these are at either end of a diametral plane through the cylindrical tablet. The machine applies force to compress the tablet until the tablet fractures. The testing machine measures the applied force (F), and also the displacement (x) of 10 the platens towards each other as the tablet is compressed. The distance (y) between the platens before force is applied, which is the diameter of the tablet, is also known. The maximum force applied is the force at 15 failure  $(F_f)$ . From this measurement of force a test parameter called diametral fracture stress, can be calculated using the equation

$$\sigma = \frac{2F_f}{\pi Dt}$$

where  $\sigma$  is the diametral fracture stress in Pascals,  $F_f$  is the applied force in Newtons to cause fracture, D is the tablet diameter in metres and t is the tablet thickness in metres.

The speed of disintegration of tablets was tested by means of a procedure in which a tablet was placed on a plastic sieve with 2mm mesh size which is immersed in 9 litres of

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demineralised water at ambient temperature of 20°C. The water conductivity is monitored until it reached a constant value. The time for dissolution of the tablets is taken as the time  $(T_{90})$  for change in the water conductivity to reach 90% of its final magnitude.

The results obtained are set out in the following table in which "comp." denotes the comparative tablets without PEG.

10	Compaction force applied(kN)	F <sub>f</sub> (Nev	vtons)	DFS (kPa	a)	T <sub>90</sub> (mi	.nutes)
		comp.	with PEG	comp.	with PEG	comp.	with PEG
	0					1.25	1.4
15	4	15.6	13.5	9.5	8.1	2.1	1.85
	9	36.5	34.8	24.5	23.6	2.5	3.0
	14	53.1	55.3	37.1	38.6	3.35	3.5

In the above table, zero compaction force denotes the particulate formulation prior to compaction.

It can be seen from this table that the presence of the PEG 1500 has very little effect on the tablet properties. In a modification to this example, the sodium acetate trihydrate used in making the base powder is also mixed with 2% of its own weight of PEG 1500.

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# Example 6

The procedure of the previous example was repeated using sodium acetate trihydrate which was mixed before use with 1% or 2% of its own weight of zeolite A24. This zeolite was as described in Example 2. Comparative tablets were made using sodium acetate trihydrate which had not been mixed with other material before use. The following results were obtained:

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Compaction force applied(kN)	F <sub>f</sub>	(Newton	ıs)	T <sub>9</sub>	(minute	es)
	zeolit	e percen	tage	zeolite	percent	age
	none	1%	2%	none	1%	2%
0				1.5	1.45	1.3
3.9	21.2	14.6	13.2	2.1	2.1	1.8
8.3	43.5	35.8	33.9	3.4	2.85	3.0
13.3	61.6	57.7	45.3	5.45	4.15	4.2

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It can be seen that here again the use of a small percentage of zeolite on the sodium acetate trihydrate to prevent caking does not have a serious deleterious effect on the tablet properties. The incorporation of sodium acetate trihydrate leads to a considerable reduction in the time for tablet dissolution, compared to tablets which do not include this material, and this benefit is also obtained when the sodium acetate trihydrate is treated beforehand with particles of zeolite as in this example or particles of PEG 1500 as in the preceding example.

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#### CLAIMS:

- A process for the production of a tablet of a compacted particulate composition by mixing
- 5 (i) a water-softening agent, and
- (ii) a water-soluble crystalline salt selected from anhydrous or hydrated sodium citrate, sodium acetate, potassium acetate and mixtures thereof and compacting the resulting mixed composition into 10 tablets or regions of tablets, characterised by the presence of particles of another substance at the surface of the crystals of the said crystalline salt (ii) before it is mixed with the water softening agent (i).
- 2. A process according to claim 1 wherein the watersoluble crystalline salt is selected from sodium citrate dihydrate, sodium acetate trihydrate, potassium acetate and mixtures thereof.
- 20 3. A process according to claim 1 or claim 2 which includes a step of applying particles of material to the surface of said crystals of the crystalline salt (ii) before the salt is mixed with other ingredients of the composition.

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4. A process for the production of a tablet of a compacted particulate composition by applying particles of material to the surface of crystals of a water-soluble

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crystalline salt selected from anhydrous or hydrated sodium citrate, sodium acetate, potassium acetate and mixtures thereof; thereafter mixing a water-softening agent and optionally other materials with said crystalline salt and compacting the resulting mixed composition into tablets or regions of tablets.

- 5. A process according to claim 4 wherein the water-soluble crystalline salt is selected from sodium citrate dihydrate, sodium acetate trihydrate, potassium acetate and mixtures thereof.
- 6. A process according to any one of the preceding claims wherein the mixed composition and the tablets or tablet regions compacted therefrom contain from 15% to 93% by weight of a water-insoluble water-softening agent.
  - 7. A process according to any one of the preceding claims wherein the mixed composition and the tablets or tablet regions compacted therefrom contain 5% to 35% by weight of sodium acetate trihydrate.
- 8. A process according to any one of the preceding claims wherein the mixed composition and the tablet or tablet regions compacted therefrom incorporate separate particles, which include detergent-active and separate particles containing at least some of said water-soluble crystalline salt.

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9. A process according to any one of the preceding claims wherein the mixed composition and the tablets or tablet regions compacted therefrom contains from 15% to 60% by weight of water-insoluble water-softening agent together with 5% to 50% by weight of one or more detergent-active compounds and from 5% to 35% by weight of sodium acetate trihydrate or potassium acetate.

10. A process according to claim 9 wherein the detergent10 active is present in particles containing water-softening agent, and the mixed composition and the tablets or tablet regions contain at least 10% and preferably at least 13% by weight of sodium acetate trihydrate with mean particle size over  $250\mu m$  preferably over  $300\mu m$ .

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11. A process according to any one of the preceding claims wherein the water-softening agent is alkali metal aluminosilicate, crystalline layered silicate or a mixture thereof.

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12. A tablet of compacted particulate composition containing a water-softening agent mixed with a crystalline salt selected from anhydrous or hydrated sodium citrate, sodium acetate, potassium acetate and mixtures thereof, characterised by particles of another material at the surface of the crystals of the said crystalline salt.

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13. A tablet according to claim 12 wherein the crystalline salt is selected from sodium citrate dihydrate, sodium acetate trihydrate, potassium acetate and mixtures thereof.

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- 14. A tablet according to claim 12 wherein said crystalline salt is sodium acetate trihydrate.
- 15. Use of crystals or anhydrous or hydrated sodium

  10 acetate, potassium acetate or mixtures of them, bearing particles of another substance at the surface of the crystals of the said salt, in a tablet of compacted particulate composition or a region thereof, to enhance the disintegration of the tablet in water.

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16. Use of crystals or anhydrous or hydrated sodium acetate trihydrate, potassium acetate or mixture of them, bearing particles of another substance at the surface of the crystals of the said salt, as an ingredient for mixing with a water-softening agent and optionally other materials to form a mixed particulate composition for compaction into a tablet of or a region thereof, wherein said crystalline salt serves to enhance the disintegration of the tablet in water.

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17. Use according to claim 15 or claim 16 wherein the said salt is sodium acetate trihydrate.





inte onal Application No PCT/EP 99/02179

A. CLASSI IPC 6	FICATION OF SUBJECT MATTER C11D17/00 C11D3/20 C11D3/12		
According to	nternational Patent Classification (IPC) or to both national classification	ation and IPC	
B. FIELDS	SEARCHED		
Minimum do IPC 6	ocumentation searched (classification system followed by classification C11D	on symbols)	
Documentat	tion searched other than minimum documentation to the extent that si	uch documents are included in the fields se	arched
Electronic d	ata base consulted during the international search (name of data bas	and, where practical, search terms used)	·
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT	<del></del>	<del></del>
Category '	Citation of document, with indication, where appropriate, of the rele	evant passages	Relevant to claim No.
P,A	EP 0 838 519 A (UNILEVER PLC ;UNI (NL)) 29 April 1998 (1998-04-29) cited in the application claims	LEVER NV	1-17
A	DE 196 37 606 A (HENKEL KGAA) 26 March 1998 (1998-03-26) page 2, line 57 - page 3, line 1 claims; example 1	.0;	1-5,7, 12-15,17
A	EP 0 002 293 A (PROCTER & GAMBLE) 13 June 1979 (1979-06-13) page 4, line 15 - page 6, line 3		1-5, 12-14
A	EP 0 711 827 A (UNILEVER PLC ;UNI (NL)) 15 May 1996 (1996-05-15) cited in the application page 4, line 6 - line 32; claims		1-6
Furt	her documents are listed in the continuation of box C.	X Patent family members are listed in	in annex.
"A" docume consider in the consider of the consideration of t	ent defining the general state of the art which is not dered to be of particular relevance document but published on or after the international date ent which may throw doubts on priority claim(s) or is cited to establish the publication date of another	"T" later document published after the inter- or priority date and not in conflict with cited to understand the principle or the invention  "X" document of particular relevance; the ci- cannot be considered novel or cannot involve an inventive step when the doc  "Y" document of particular relevance; the ci-	the application but laimed invention be considered to cument is taken alone
"O" docum other "P" docum	in or other special reason (as specified)  ent referring to an oral disclosure, use, exhibition or  means  ent published prior to the international filing date but  han the priority date claimed	cannot be considered to involve an inv document is combined with one or mo ments, such combination being obviou in the art.  "&" document member of the same patent if	re other such docu- us to a person skilled
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PCT/EP	99/02179	)
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Patent document cited in search report		Publication date		atent family nember(s)	Publication date
EP 0838519 A		29-04-1998	GB	2318575 A	29-04-1998
DE 19637606	Α	26-03-1998	NONE		
EP 0002293	А	13-06-1979	BE DE IT NL NL CA FR GB US	56 T 2857472 A 1101296 B 7815064 A 7815064 T 1109752 A 2443502 A 2041966 A 4219435 A	18-04-1980 17-07-1980 28-09-1985 30-05-1980 30-05-1980 29-09-1981 04-07-1980 17-09-1980 26-08-1980
EP 0711827	 А	15-05-1996	NONE		

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